

from DHHS should be directed to Andrew J. Hagan, OCSE (telephone (202) 401-5375). This is not a toll-free number. Any written comments on the minutes should be directed to Ms. Samara Weinstein, Executive Director of the Working Group, as shown above.

Signed at Washington, DC, this 29th day of March, 1999.

**David Gray Ross,**

*Commissioner, Office of Child Support Enforcement.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. 99N-0038]

#### Agency Information Collection Activities; Announcement of OMB Approval; Survey of Biomedical Equipment Manufacturers for Year 2000-Compliant Products

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing that a collection of information entitled "Survey of Biomedical Equipment Manufacturers for Year 2000-Compliant Products" has been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

**FOR FURTHER INFORMATION CONTACT:** Peggy Schlosburg, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1223.

**SUPPLEMENTARY INFORMATION:** In the **Federal Register** of January 22, 1999 (64 FR 3524), the agency announced that the proposed information collection had been submitted to OMB for review and clearance under 44 U.S.C. 3507. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. OMB has now approved the information collection and has assigned OMB control number 0910-0397. The approval expires on September 30,

1999. A copy of the supporting statement for this information collection is available on the Internet at "http://www.fda.gov/ohrms/dockets".

Dated: March 29, 1999.

**William K. Hubbard,**

*Acting Deputy Commissioner for Policy.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

#### Public Input on Public Health; Open Public Forum

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice of meetings.

**SUMMARY:** The Food and Drug Administration (FDA), Office of Consumer Affairs and Office of Regulatory Affairs, Pacific Region, is announcing a series of open public forums entitled: "Public Input on Public Health, FDA Listens to You, A Town Hall Meeting." The purpose of the forums is to provide an opportunity for FDA's primary stakeholders, U.S. consumers, to have an open dialogue with FDA's senior policy makers about their consumer protection concerns. FDA plans to use the information in the development of the Pacific Region Strategic Plan and in the development of FDA's nation wide priorities. Under the FDA Modernization Act of 1997 (FDAMA), FDA was mandated by Congress to have ongoing consultations with its stakeholders on how FDA can best meet their regulatory requirements and to protect the public health. Two issues of particular concern, this year, are strengthening the science base of the agency and improving risk-based communication with the public.

**DATES:** Send registration and requests for oral presentations by May 5, 1999. See Table 1 in section II of this document for a complete schedule of all the meetings.

**ADDRESSES:** Send written comments to the specific contact person. See Table 1 in section II of this document for a complete listing of meeting locations and contact persons.

**FOR FURTHER INFORMATION CONTACT:**

For general information: James Rowell

or Patricia Alexander, Food and Drug Administration, 5600 Fishers Lane, rm. 16-75, Rockville, MD 20857, 301-827-4414 or 301-827-4391, FAX 301-443-9767.

For specific meeting information: See Table 1 in the **SUPPLEMENTARY INFORMATION** section of this document.

#### SUPPLEMENTARY INFORMATION:

##### I. Background

FDA is a science based consumer protection agency responsible for ensuring that: (1) Foods are safe, wholesome and sanitary; (2) human and veterinary drugs, biological products, and medical devices are safe and effective; (3) cosmetics and electronic products that emit radiation are safe. FDA also ensures that these regulated products are honestly and accurately labeled and in compliance with applicable laws and regulations. FDA strives to maximize public health protection while reducing regulatory burdens.

Public participation in these forums will provide an essential ingredient to the achievement of the Pacific Region's intermediate and long range strategic planning goals. Additional benefits include: (1) Providing the opportunity to hear directly from consumers their concerns about health and policy issues, (2) reaching out to a broad representation of community based and consumer organizations to reach the full diversity of consumers, (3) using the information gained at these forums in FDA's decisionmaking process, (4) obtaining information necessary for the development of innovative programs to raise public awareness, (5) fostering communication among local agencies, both public and private, in order to more effectively respond to the public's need for information that empowers them in making health related decisions, and (6) encouraging individuals to take personal responsibility for protecting their own health.

##### II. Scheduled Meetings

The open public forums will be held in several locations throughout the country. The scheduled date and time, location, and specific contact person for each meeting is listed in Table 1 as follows:

TABLE 1.—MEETING SCHEDULES AND CONTACTS FOR REGISTRATION

Date	Time	Place	Address	Contact
Wednesday, May 12, 1999	10 a.m. to 1 p.m.	Elihu Harris State Office Bldg. Auditorium.	1515 Clay St., Oakland, CA.	Mary Ellen Taylor at 510-337-6888, FAX 510-337-6708